

Europäisches Patentamt

European Patent Office

Office européen des brevets



(11) EP 0 769 551 A1

(12)

EUROPEAN PATENT APPLICATION

published in accordance with Art. 158(3) EPC

- (43) Date of publication: 23.04.1997 Bulletin 1997/17
- (21) Application number: 96905037.6
- (22) Date of filing: 08.03.1996

- (51) Int. Cl.⁶: C12N 15/00, C12N 9/90
- (86) International application number: PCT/JP96/00574
- (87) International publication number:WO 96/28545 (19.09.1996 Gazette 1996/42)
- (84) Designated Contracting States:
 AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC
 NL PT SE
- (30) Priority: 10.03.1995 JP 51234/95
- (71) Applicant: KIRIN BEER KABUSHIKI KAISHA Chuo-Ku, Tokyo 104 (JP)
- (72) Inventors:
 - KAJIWARA, Susumu Tokyo 158 (JP)

- MISAWA, Norihiko, Kirin Beer KK Kanazawa-ku, Yokoham-a-shi, Kanagawa 236 (JP)
- KONDO, Keiji, Kirin Beer KK Kanazawa-ku, Yokohama-shi, Kanagawa 236 (JP)
- (74) Representative: Kolb, Helga, Dr. Dipl.-Chem. et al Hoffmann, Eitle & Partner, Patent-und Rechtsanwälte, Arabellastrasse 4 81925 München (DE)

(54) DNA STRAND USEFUL IN INCREASING CAROTENOID YIELD

(57) A DNA chain having characteristic of increasing carotenoid production, and containing the nucleotide sequence which encodes the polypeptide having the substantially same amino acid sequence described in SEQUENCE ID No. 1 or 2, or a DNA chain which hybridizes with said DNA chain, and a method for production for carotenoid characterized by introducing said DNA chain into the carotenoid-producing microorganisms, culturing said transformed microorganism and increasing carotenoid content in the culture broth and cells.

Description

10

FIELD OF THE INVENTION

The present invention relates to a DNA chain which provides higher carotenoid content during biosynthesis of carotenoid and a method for producing carotenoids characterized by introducing said DNA chain into carotenoid producing microorganism to express said chain and to obtain higher carotenoid content.

BACK GROUND OF THE INVENTION

Carotenoid is a general name of a kind of natural pigments. Generally, carotenoids have 40 carbon atoms and consists of isoprene skeletons, and Carotenoids are abundant in the natural world. Approximately 600 kinds of carotenoids have been isolated and identified up to the present [(see Key to carotenoids. Basel-Boston, Birkhauser, 1987(Pfander, H. ed.)]. Carotenoids are synthesized through the isoprenoid biosynthetic pathway, a part of which is common to the pathways for steroids and other terpenoids. Passing through the isoprene common biosynthetic pathway, hydroxymethylglutaryl-CoA(HMG-CoA) is converted to isopentenyl pyrophosphate(IPP), which has 5 carbon atoms, via mevalonate. Then IPP is converted to dimethylallyl pyrophosphate(DMAPP) by isomerization. Then, by polycondensation with IPP which has 5 carbon atoms, DMAPP is converted sequentially to geranyl pyrophosphate(GPP which has 10 carbon atoms), farnesyl pyrophosphate (FPP which has 15 carbon atoms), geranylgeranyl pyrophosphate(GGPP which has 20 carbon atoms) and so forth (Figure 1).

The carotenoid biosynthetic pathway is branched from the isoprene common pathway at the point of GGPP is formed. At the point, two molecules of GGPP are condensed to synthesize phytoene which is the first carotenoid and colorless. Then, phytoene is converted to lycopene by desaturation reaction. Then, lycopene is converted to β -carotene by cyclization. Various xanthophylls such as zeaxanthin and astaxanthin are synthesized by introducing hydroxyl groups or keto groups to β -carotene.

Recently, the inventors of the present invention cloned the carotenoid biosynthesis genes derived from Erwinia uredovora, which is a non-photosynthetic epiphytic bacterium in Escherichia coli by using yellowish color of Er. uredovora as markers and elucidated the functions of the genes. Then, various combinations of these genes are introduced to express, and it made possible that microorganisms such as <u>E. coli</u> and yeast produce phytoene, lycopene, β-carotene, zeaxanthin and so forth(See Figure 2): [See Misawa, N., Nakagawa, M., Kobayashi, K., Yamano, S., Izawa, Y., Nakamura, K. and Harashima, K., "Elucidation of the Erwinia uredovora carotenoid biosynthetic pathway by functional analysis of gene products expressed in Escherichia coli", J. Bacteriol., 172: 6704-6712 (1990); Misawa, N., Yamano, S., and Ikenaga, H., "Production of β-carotene in Zymomonas mobilis and Agrobacterium tumefaciens by introduction of the biosynthesis genes from Erwinia uredovora", Appl. Environ. Microbiol., 57: 1847-1849 (1991); Yamano, S., Ishii, T., 35 Nakagawa, M., Ikenaga, H., and Misawa, N., "Metabolic engineering for production of β-carotene and lycopene in Saccharomyces cerevisiae", Biosci. Biotech. Biochem., 58: 1112-1114 (1994) and Japanese Patent Application laid-open No. HEI 3-58786(Japanese Patent Application filing No. HEI 2-53255): "A DNA chain useful for synthesis of carotenoids" by the inventors of the present invention]. With the carotenoid biosynthesis genes from Er. uredovora, carotenoids can be synthesized from FPP. Since FPP is the common substrate not only for carotenoids but also for steroids and other terpenoids, bacteria incapable of synthesizing carotenoids also have FPP. Accordingly, for example, when four crt genes, crtE, crtB, crtI and crtY, which are necessary for biosynthesis of β-carotene from FPP are introduced in microorganisms, the microorganism becomes capable of producing β-carotene (See Figure 2). Furthermore, by the same procedures as mentioned above, the inventors cloned the carotenoid biosynthesis genes derived from a marine bacterium, Agrobacterium aurantiacum in E. coli. By expressing various combinations of the genes from the bacterium and those from the above-mentioned Er. uredovora, it made possible that the microorganisms such as E. coli produce astaxanthin, canthaxanthin and so forth (See Figure 3): (Norihiko Misawa et al., "Elucidation of an astaxanthin biosynthetic pathway at the level of the biosynthesis genes", Abstract of the 36th Symposium on the chemistry of natural products: 175-180 (1994)). Among the above carotenoids, astaxanthin, zeaxanthin and β-carotene are already in practical use and are regarded as promising substances. They are used for food or feed additives as red or yellow natural coloring agents or as nutritional aid having cancer prophylactic activity, immunopotentiating activity or provitamin A activity. Accordingly, when the carotenoid biosynthesis genes obtained by the inventors is used as exogenous genes for transforming microorganisms such as E. coli to express, it gave microorganisms such as E. coli the capability of biosynthesis for producing useful carotenoids. Up to now, it is the only way to improve production of useful carotenoids was to find out microorganism which can synthesize sufficient amount of a targeted carotenoid, and to try to increase its production by investigating culture conditions or mutation treatment. Owing to the studies done by the inventors, it became possible to choose host microorganism which is cultured easily and proliferates rapidly, and is guaranteed to be safe for food regardless of its carotenoid producing capability. As a matter of course, it is also possible to use microorganisms which can synthesize sufficient amount of useful carotenoids originally. In such a case, by transforming the microorganisms with carotenoid biosynthesis genes, it became possible to obtain higher carotenoid production or to alter final caroten-

oid products. For example, when both $\underline{crt}W$ and $\underline{crt}Z$ genes from \underline{Ag} , $\underline{aurantiacum}$ were introduced into a microorganism capable of producing β -carotene as a final product to express them, the microorganism was transformed to another one which produce astaxanthin as a final product.

On the other hand, both astaxanthin and β-carotene can also be synthesized by organic synthesis methods. In these cases, considering these carotenoids are used for feed or food additives, there is problems that by-products are also produced and such synthetic products are not preferred by consumers because they prefer natural products. However, carotenoids produced by the conventional fermentation methods could not compete with those by the organic synthesis methods in price. As mentioned earlier, when the above mentioned carotenoid biosynthesis genes are used, it improves the fermentation methods, thereby it is considered that the carotenoid produced by the fermentation methods will be able to compete with those by the organic synthesis methods in price. If the microorganism can accumulate enough amount of carotenoid in itself, the carotenoid produced by the microorganisms will succeed in such price competition. Therefore, a technology to obtain higher carotenoid content by using microorganisms has been longed for.

Until now, in order to obtain higher carotenoid production in its biosynthesis; the traditional random mutation method is only employed to select mutant strains having higher carotenoid content with mutagenic agent such as NTG. However, this method requires huge amount of time and labor of technicians. In addition, even if enhancement of carotenoid synthesis is successfully achieved, the method requires both huge amount of time and effort to inhibit decreasing of carotenoid content caused by frequent reverse mutations naturally happens because the method lacks its theoretical basis.

SUMMARY OF THE INVENTION

The object of the present invention is to increase amount of carotenoids biosynthetically produced by microorganisms.

To solve the above problem, the inventors have investigated the problem thoroughly and developed a novel technology which provides several times higher carotenoid production amount by introducing a DNA chain containing only one gene into a carotenoid producing microorganism to express the gene in them.

More specifically, the inventors of the present invention found the followings and completed the present invention. When a DNA chain containing a gene substantially encoding an amino acid sequence of IPP isomerase which converts IPP into DMAPP, is introduced in microorganisms such as <u>E. coli</u> having carotenoid synthesis gene derived from <u>Er. uredovora</u> and so forth, content of carotenoid in cells such as lycopene and β-carotene becomes 1.5-4.5 times higher than that in control cells can be achieved. The gene substantially encoding IPP isomerase amino acid sequence which converts IPP into DMAPP was obtained from the astaxanthin producing microorganisms such as <u>Phaffia rhodozyma</u> and <u>Haematococcus pluvialis</u>.

The characteristics of the DNA chain of the present invention are as follows.

- (1) A DNA chain capable of increasing carotenoid production amount and containing the nucleotide sequence which encodes the polypeptide having the amino acid sequence substantially shown in Sequence ID No. 1, or a DNA chain that can be hybridized with said DNA chain.
- (2) A DNA chain capable of increasing carotenoid production and containing the nucleotide sequence which encodes the polypeptide having the amino acid sequence substantially shown in Sequence ID No. 2, or a DNA chain that can be hybridized with said DNA chain.

The present invention also relates to a method for carotenoid production. The characteristics of the carotenoids production methods of the present invention are as follows.

- (3) A production method characterized by introducing the DNA chain mentioned above either (1) or (2) into carotenoid producing microorganism, culturing said transformed microorganism and increasing carotenoid content in the cells and culture broth.
- (4) A production method characterized by introducing the DNA chain containing the nucleotide sequence which encodes the polypeptide having the substantially same amino acid sequence shown in Sequence ID No. 3, or a DNA chain that can be hybridized with said DNA chain into carotenoid producing microorganism, culturing said microorganism and increasing carotenoid content in the cells and culture broth.

The present invention is described herein below.

As described in before, by introducing the carotenoid biosynthesis gene derived from microorganisms such as Erwinia uredovora, the non-photosynthetic soil bacteria and Agrobacterium aurantiacum, the marine bacteria) into other microorganisms which do not produce carotenoids such as E. coli, the microorganism can produce useful carotenoids such as astaxanthin, zeaxanthin, β-carotene and lycopene. In order to compete in price of the carotenoid produced by using the organic synthesis methods, it is necessary to achieve as higher carotenoid production as possible. The IPP isomerase gene, which include the gene encoding the polypeptide whose amino acid sequence is substantially IPP isomerase, of the invention is extremely useful for increasing the production amount of carotenoids. By using mod-

35

40

45

ern biotechnology, it is relatively easy to increase production amount of a protein encoded by an exogenous gene by enhancing expression level of the gene. However, if amounts of substrate necessary for a protein, that is enzyme, is limited, higher production of the protein does not lead to higher production of biochemicals such as carotenoids. For example, without sufficient amount of FPP, which is the first substrate, enhancement of expression level of the carotenoid synthesis genes does not lead to higher amount of carotenoids production. This time, we succeeded in increasing carotenoid production amount by introducing the IPP isomerase gene. It is considered that the introduction of the IPP isomerase gene makes the flow of, the upstream of the pathway up to FPP larger(more efficient) and consequently, increased supply of FPP led to higher carotenoid production amount. The present invention started from the findings that by introducing either the gene encoding IPP isomerase, which convert from IPP to DMAPP vise versa, or encoding the protein homologous to IPP isomerase into carotenoid producing microorganism such as E. coli, to express the gene, carotenoid production amount is increased. By using carotenoid biosynthesis genes from Er. <u>uredovora,</u> cDNA expression libraries of Phaffia rhodozyma. Haematococcus pluvialis and so forth were prepared in β-carotene producing \underline{E} , coli as a host. As increased β -carotene content in \underline{E} , coli made, some of the yellowish colonies brighter till almost orange. The plasmids extracted from such E. coli colonies were analyzed and were found to have genes with high homology to IPP isomerase of Saccharomyces cerevisiae. It has been speculated that HMG-CoA reductase(Figure 1), which catalyzes the reaction from HMG-CoA to mevalonate, may be the rate limiting enzyme for terpenoids including carotenoids. However, as for IPP isomerase, any such report has not been presented. Therefore, increase of carotenoid production by introducing a IPP isomerase gene was a new finding.

The present invention provides a DNA chain having characteristics of increasing carotenoid production amount, and it containing the nucleotide sequence which encodes the polypeptide having the substantially same amino aid sequence as those of IPP isomerase, and a production method for carotenoid characterized by introducing said DNA chain into the carotenoid producing microorganism, culturing said transformed microorganism and increasing carotenoid content in the culture broth and cells.

The DNA chains of the present invention includes the DNA chains mentioned above (1) or (2), or the DNA chains which hybridize to said chains under stringent conditions.

Substantially, the polypeptides encoded by the DNA chains of the present invention have the amino acid sequences shown in SEQUENCE ID No. 1(A-B in Figures 4 and 5) or in SEQUENCE ID No. 2(C-D, in Figures 6 and 7). In the present invention, the polypeptides encoded by these DNA chains, the proteins of which amino acid sequence is substantially IPP isomerase, may be altered by deletion, replacement, addition and so forth of some amino acids, as long as the resulted polypeptides hold their higher carotenoid production activity. This allowance corresponds to "having the substantially same amino acid sequence substantially shown in SEQUENCE ID No. 1 or No. 2". As an example, a sequence which lacks the first amino acid(Met) can be included as the altered polypeptide or the altered enzyme. Needless to say, the DNA chains of the present invention include not only the chains having the nucleotide sequences which encode the amino acid sequences shown in SEQUENCE ID No. 1 and 2(Figures 4 to 5), but also the degenerate isomers of the chains, which differs only on degenerate codons and encode the same polypeptides as the original chains do.

(1) Obtaining the DNA chains

40

One method to obtain a DNA chain having the nucleotide sequence which encodes the amino acid sequence of the above protein is chemical synthesis of the DNA chain at least a part of the chain according to the known nucleic acid synthesis method. However, considering that there are so many amino acids bound in the protein, it would be more preferable than chemical synthesis to make cDNA libraries of Haematococcus pluvialis or Phaffia rhodozyma or the like to obtain a targeted DNA chain by applying some popular method in the field of genetic engineering such as hybridization with appropriate probes.

(2) Transformation of microorganisms such as E. coli and expression of gene

Higher carotenoid content in culture broth or cells of microorganisms can be achieved by introducing the above mentioned DNA chain of the present invention into appropriate microorganisms such as carotenoid-producing bacteria such as <u>E. coli</u> and <u>Zymomonas mobilis</u> containing carotenoid biosynthesis genes from <u>Erwinia uredovora</u> and so forth, or carotenoid-producing yeast such as <u>Saccharomyces</u> <u>cerevisiae</u> containing carotenoid biosynthesis genes from <u>Erwinia uredovora</u> and so force.

The outline of the method to introduce exogenous genes into preferable microorganisms is mentioned below.

Procedures or methods to introduce and express exogenous genes in microorganisms such as <u>E. coli</u>, besides those mentioned below in the present invention, includes those widely used in the field of genetic engineering. Those are applicable to the invention. See "Vectors for cloning genes", Methods in Enzymology, 216: 469-631 (1992), Academic Press; "Other bacterial systems", Methods in Enzymology, 204: 305-636 (1991) Academic Press).

[E. coli]

There are some established and efficient methods to introduce exogenous genes to <u>E. coli</u> such as Hanahan's method and rubidium method, and they are applicable to the present invention (See Sambrook, J., Fritsch, E. F., Maniatis, T., "Molecular cloning-A laboratory manual", Cold Spring Harbor Laboratory Press (1989)). Expression of exogenous genes in <u>E. coli</u> can be performed by known methods (See "Molecular cloning-A laboratory manual", ibid.), for example, vectors for <u>E. coli</u> such as pUC and pBluescript vectors having <u>lac</u> promoter can be used. The inventors of the present invention used pSPORT1 vector or pBluescript II KS vector having <u>lac</u> promoter as vectors for <u>E. coli</u>, and inserted the IPP isomerase gene, derived from <u>Haematococcus pluvialis</u>, <u>Phaffia rhodozyma</u> or <u>Saccharomyces cerevisiae</u>, into the <u>lac</u> promoter with the direction of reading through of the transcription, and expressed the gene in <u>E. coli</u>.

[Yeast]

There are some established methods such as the lithium method to introduce exogenous genes into <u>Saccharomy-ces cerevisiae</u>, yeast, and such methods are applicable to the present invention (See "New biotechnology on yeast", Ed. Bio-industry Association (Yuichi Akiyama, editor in chief), Igaku Syuppan Center). Expression of exogenous genes in yeast can be performed as follows. Using both promoters and terminators, e.g. for <u>PGK</u> and <u>GPD</u>, an expression cassette is constructed by inserting the exogenous gene so that during transcription, the gene is to be read through at the position between the promoter and the terminator. Expression can be performed by inserting the expression cassette into a vector for <u>S. cerevisiae</u> such as YRp vectors (multi-copy vectors for yeast, replication starts at ARS sequence of yeast chromosome), YEp vectors (multi-copy vectors for yeast, replication starts at 2μm DNA) and Ylp vectors (vectors for yeast chromosome, no starting point of replication in yeast) (See "New biotechnology on yeast", ibid.; "Genetic engineering for production of substances", Ed. Japanese Society of Agrocultural Chemistry, Asakura Publishing company; or Yamano, S., Ishii, T., Nakagawa, M., Ikenaga, H., Misawa, N., "Metabolic engineering for production of β-carotene and lycopene in <u>Saccharomyces cerevisiae</u>", Biosci. Biotech, Biochem., 58: 1112-1114 (1994)).

[Zymomonas mobilis]

Introduction of exogenous genes into Zymomonas mobilis, the ethanol-producing bacterium can be performed by conjugal transfer method which is commonly used for gram negative bacteria. Expression of exogenous gene in Zymomonas mobilis can be performed by using pZA22 vector for this bacterium (See Katsumi Nakamura, "Molecular breeding of Zymomonas bacteria", Journal of the Japanese Society of Agrocultural Chemistry, 63: 1016-1018 (1989); and Misawa, N., Yamano, S., Ikenaga, H., "Production of β-carotene in Zymomonas mobilis and Agrobacterium tume-faciens by introduction of the biosynthesis genes from Erwinia uredovora", Appl. Environ. Microbiol., 57: 1847-1849 (1991)).

(3) Method to increase carotenoid production in microorganisms

By applying the above mentioned procedures or methods for introduction and expression of exogenous genes in microorganisms, both the carotenoid synthesis genes and the IPP isomerase gene can be introduced to express, and microorganisms capable of producing large amount of carotenoid can be obtained.

Farnesyl pyrophosphate (FPP) is the common substrate not only for carotenoids but also for other terpenoids such as sesquiterpenes, triterpenes, sterols and hopanols. In general, since microorganisms are synthesizing terpenoids even though they are not capable of synthesizing carotenoids, basically all of the microorganisms possesses FPP as 45 an intermediate metabolite. On the other hand, Erwinia uredovora, the non-photosynthetic bacterium having the carotenoid synthesis genes can synthesize up to several useful carotenoids such as lycopene, β-carotene, zeaxanthin by using FPP as a substrate. When the genes are combined with the carotenoid synthesis genes of Agrobacterium aurantiacum, the marine bacterium, up to several useful carotenoids such as cantaxanthin and astaxanthin can also be synthesized (See Figures 2 and 3). The inventors of the present invention already confirmed that by introducing crt genes 50 of Erwinia uredovora into microorganisms such as Saccharomyces cerevisiae, yeast and Zymomonas mobilis, ethanolproducing bacteria; these microorganisms can produce carotenoids such as β-carotene as anticipated [Yamano, S., Ishii, T., Nakagawa, M., Ikenaga, H., Misawa, N., "Metabolic engineering for production of β-carotene and lycopene in Saccharomyces cerevisiae", Biosci. Biotech, Biochem., 58:1112-1114 (1994); Misawa, N., Yamano, S., Ikenaga, H., "Production of β-carotene in Zymomonas mobilis and Agrobacterium tumefaciens by introduction of the biosynthesis 55 genes from Erwinia uredovora", Appl. Environ. Microbiol., 57:1847-1849 (1991); and Japanese laid-open Patent Application No. HEI 3-58786(Japanese Patent Application filing No. HEI 2-53255):"A DNA chain useful for synthesis of carotenoids" by the inventors].

From the above findings, it can be expected that when an appropriate combinations of the carotenoid synthesis genes derived from <u>Er. uredovora</u> and those from marine bacteria(typically the carotenoid synthesis genes derived from

Ag. aurantiacum) are introduced into the same microorganism simultaneously, as a principle, all of the microorganisms, in which such genes are introduced and of which introduction-expression system is established, can produce useful carotenoids such as astaxanthin and zeaxanthin.

In such cases, if the IPP isomerase gene(typically, derived from <u>Haematococcus pluvialis</u>, <u>Phaffia modozyma</u> and <u>Saccharomyces cerevisiae</u>) is introduced according to the above mentioned method, and is expressed concomitantly with the above carotenoid synthesis gene, higher production amount of useful carotenoids can be achieved.

(4) Deposit of the microorganisms

The recombinant E. coli strain JM109 has been deposited as follows with the National Institute of Bioscience and Human-Technology, the Agency of Industrial Science and Technology. The strain contains the plasmid having the isolated gene which is the DNA chain of the invention. The names of the plasmids are shown in the parentheses.

(i) JM109(pRH1)

Deposit No.: FERM BP-5032 Date of Receipt: March 6th, 1995

(ii) JM109(pHP11)

Deposit No.: FERM BP-5031
Date of Receipt: March 6th, 1995

(ii) JM109(pSl1)

Deposit No.: FERM BP-5033 Date of Receipt: March 6th, 1995

BRIEF DESCRIPTION OF THE DRAWINGS

25

45

10

15

20

FIGURE 1 shows the isoprene common biosynthetic pathway from HMG-CoA to FPP.

FIGURE 2 shows the carotenoid biosynthetic pathway, and the functions of the carotenoid synthesis genes of <u>Erwinia uredovora</u>, the non-photosynthetic bacterium.

FIGURE 3 shows the carotenoid biosynthetic pathway, and the functions of the carotenoid synthesis genes of <u>Agrobacterium aurantiacum</u>, the marine bacterium. The solid line shows major biosynthetic pathway and the dotted line shows minor one.

FIGURES 4 and 5 shows the nucleotide sequence of the IPP isomerase gene and the amino acid sequence of the polypeptide encoded by said gene of <u>Phaffia rhodozyma</u>, the astaxanthin-producing yeast. In the Figure, the sequence from mark A to B shows the open reading frame encoding the polypeptide consisting of 251 amino acids.

FIGURES 6 and 7 shows the nucleotide sequence of the IPP isomerase gene and the amino acid sequence of the polypeptide encoded by said gene of <u>Haematococcus pluvialis</u>, the astaxanthin-producing green alga. In the Figure, the sequence from mark C to D shows the open reading frame encoding the polypeptide consisting of 259 amino acids.

FIGURES 8 and 9 shows the nucleotide sequence of the IPP isomerase gene and the amino acid sequence of the polypeptide encoded by said gene of <u>Saccharomyces cerevisiae</u>, the yeast for laboratory use. In the Figure, the sequence from mark E to F shows the open reading frame encoding the polypeptide consisting of 288 amino acids.

FIGURE 10 shows the plasmids containing the carotenoid biosynthesis genes of <u>Erwinia uredovora</u>, the non-photosynthetic bacterium.

FIGURE 11 shows the plasmids containing the IPP isomerase gene of <u>Phaffia rhodozyma</u>, <u>Haematococcus pluvialis</u>, or <u>Saccharomyces cerevisiae</u>.

FIGURE 12 shows the growth curve in the culture broth of the lycopene producing <u>E. coli</u> strains(L:). In the Figure, "control" means the <u>E. coli</u> strain having no exogenous IPP isomerase gene.

FIGURE 13 shows the lycopene production curve in the culture broth of the lycopene producing <u>E. coli</u> strains(L:). In the Figure, "control" means the <u>E. coli</u> strain having no exogenous IPP isomerase gene.

FIGURE 14 shows production of lycopene(L:), β -carotene(β :) and phytoene(P:) in the cultured cells of the <u>E. coli</u> strains. In the Figure, "control" means the <u>E. coli</u> strain having no exogenous IPP isomerase gene.

EXAMPLE

The following examples illustrate the present invention in more detail, however, the present invention is not limited to them. The genetic recombination experiments used here are based on the standard methods(Sambrook, J., Fritsch, E. F., Maniatis, T., "Molecular cloning-A laboratory manual", Cold Spring Harbor Laboratory Press (1989)) unless otherwise stated.

(EXAMPLE 1) Biological materials and culture conditions

Phaffia rhodozyma ATCC 24230 Strain(Astaxanthin-producing yeast) registered at the American Type Culture Collection(ATCC) is used. YM media(yeast extract 0.3%, malt extract 0.3%, bactopeptone 0.5%, Glucose 1%) is used for Ph. rhodozyma. Haematococcus pluvialis, the astaxanthin-producing green alga, NIES-144 strain registered at the Global Environmental Forum is used. Ha. pluvialis is cultured at 20°C for about 4 days in basic culture media(yeast extract 0.2%, sodium acetate 0. 12%, L-asparagin 0. 04%, magnesium chloride hexahydrate 0. 02%, ferrous sulfate heptahydrate 0.001%. calcium chloride dihydrate 0.002%) under 12 hr light (20 μΕ/m²s)/12 hr dark condition. Furthermore, in order to induce astaxanthin synthesis in Ha. pluvialis, cyst formation, a kind of differentiation, has to be induced. To induce cyst formation, both acetic acid 45 mM and ferrous sulfate heptahydrate 450 μM at final concentrations are added. Ha. pluvialis in the media is cultured for about 12 hr at 20°C with light(125 μΕ/m²s). Saccharomyces cerevisiae(Yeast for laboratory use) S288C strain registered at the Yeast Genetic Stock Center is used. For Sa. cerevisiae, YPD media(yeast extract 1%, bactopeptone 2%, glucose 2%) is used.

15 (EXAMPLE 2) Preparation of whole RNA in Phaffia rhodozyma

Phaffia rhodozyma ATCC 24230 strain is cultured with shaking for approx. 24 hr at 20°C in 400 ml of YM media. When the turbidity of the media reached at OD₆₀₀ = 0.4, the bacteria are collected and frozen in liquid nitrogen. The frozen bacteria are stored in the freezer at -80°C and used for preparing total RNA. After thawing the frozen bacteria in a tube on ice, the bacteria are suspended in 6 ml of ANE buffer(10 mM sodium acetate, 100 mM sodium chloride, 1 mM EDTA, pH 6.0). Glass beads are added to cover the surface of the bacteria layer. Then, 600 μl of 10% SDS and 6 ml of phenol prewarmed at 65°C are added. The suspension is kept at 65°C for 5 minutes, and the tube is vortexed to crushed cell membranes at every 30 seconds. Then, the suspension is rapidly cooled down to room temperature and centrifuged for 10 minutes at 1,500 x g at room temperature. Equal volume of phenol is added to the supernatant and vortex for 2 minutes. Then the suspension was centrifuged for 10 minutes at 1,500 x g at room temperature. Then, by using equal volume of phenol/chloroform(1/1(v/v)) and chloroform alone, the same procedures as above are performed. To the resulted supernatant, one tenth volume of 3 M sodium acetate and three volume of ethanol are added; then the supernatant is stored in the freezer at -20°C for 30 minutes. The supernatant is centrifuged for 15 minutes at 15,000 x g at 4°C, a pellet is rinsed with 70% ethanol and dried. The residual is dissolved in 200 μl of sterilized water to make total RNA solution of Ph. rhodozyma. By this preparation procedure, 1.6 mg of total RNA is obtained.

(EXAMPLE 3) Preparation of whole RNA in Haematococcus pluvialis

Haematococcus pluvialis NIES-144 strain is cultured for approx. 4 days in 800 ml of the basic culture media under the condition of 20°C, light intensity at 20 μE/m²s and 12 hr light/12 hr dark cycle. Then, both acetic acid 45 mM and ferrous sulfate heptahydrate 450 μM as final concentrations are added. The H. pluvialis in the media is cultured for approx. 12 hr at 20°C with light (125 μE/m²s). The bacteria are collected from the media, frozen in liquid nitrogen arid crushed in the mortar to give powder. Then, three ml of ISOGEN-LS[Nippon Gene K.K.] is added to the powder and stand for 5 minutes. Then 0.8 ml of chloroform is added, and the solution is stirred vigorously for 15 seconds and stand at room temperature for 3 minutes. The solution is centrifuged for 15 minutes at 4°C, 12,000 x g, two ml of isopropanol is added to the supernatant and the supernatant is stood at room temperature for 10 minutes. Then, the solution is centrifuged for 10 minutes at 4°C, 12,000 x g. The resulted pellet is rinsed with 70% ethanol to dry. After drying, the residual is dissolved in 1 ml of TE buffer(10 mM Tris-HCl pH 8.0, 1 mM EDTA) to make total RNA solution of Ha. pluvialis. By this preparation procedure, 4.1 mg of whole RNA was obtained.

(EXAMPLE 4) Establishing cDNA expression libraries of Phaffia rhodozyma and Haematococcus pluvialis

By using Oligotex-dT30 Super[Takara Syuzo K.K.], poly A + RNA from <u>Phaffia rhodozyma</u> and <u>Haematococcus pluvialis</u> are purified from approx. 1 mg total RNA respectively. The purification is performed according to the methods mentioned in the package insert. By following the method, approx. 26 μ g of poly A + mRNA from <u>Ph. rhodozyma</u> and approx. 14 μ g of it from <u>Ha. pluvialis</u> are purified.

Preparation of cDNA is performed with Superscript™ plasmid system(GIBCO BRL) by the method mentioned in the package insert with some modifications. Approx. 5 µg of poly A + mRNA is used. A synthetic DNA consisting of the recognition sequence for the restriction enzyme Not1 and 15 mers oligo-dT is used as a primer. The complementary DNA is synthesized with reverse transcriptase, SUPERSCRIPT RT. Then, by using Escherichia coli DNA ligase, E. coli DNA polymerase and E. coli RNase H, double strand DNA is synthesized. Then, the linker of the restriction enzyme Sall is bound by using T4 DNA ligase. cDNA is designed to have the Sall site at the upstream terminal of itself and the Not1 site at the downstream of poly A. Fractionation by size of these cDNAs is performed by electrophoresis and the fractions ranging from 0.7 kb to 3.5 kb are collected. cDNA in the collected fractions is ligated to cDNA expression vector

pSPORT I NotI-SalI-Cut by using both the ligation buffer which is included in the kit, 50 mM Tris-HCl pH 7.6, 10 mM MgCl₂, 1 mM ATP, 1 mM DTT, 5% PEG 8,000 and T4 DNA Ligase. The cDNA expression vector pSPORT I has <u>lac</u> promoter at the upstream of the <u>SalI</u> site and can express cDNA in <u>E. coli</u>. Then, by using whole the ligated DNA solution, transformation of the competent cells of <u>E. coli</u> DH5α prepared is performed according to the method described in "Molecular Cloning 2nd edition: Cold Spring Harbor Laboratory, 1.21-1.41(1989). Approx. 200,000 transformed strains of <u>Ph. rhodozyma</u> and approx. 40,000 transformed strains of <u>Ha. pluvialis</u> are obtained. After collecting all of the transformants, the plasmid DNA is prepared according to the method described in "Molecular Cloning 2nd edition, ibid." As a result, 0.9 mg and 0.6 mg of plasmid DNAs are obtained respectively and these are assigned as cDNA libraries of <u>Ph. rhodozyma</u> and <u>Ha. pluvialis</u>.

(EXAMPLE 5) Preparation of carotenoid-producing E. coli

10

The plasmid pCAR16(Misawa, N., Nakagawa, M., Kobayashi, K., Yamano, S., Izawa, Y., Nakamura, K., Harashima, K., "Elucidation of the <u>Erwinia uredovora</u> carotenoid biosynthetic pathway by functional analysis of gene products expressed in <u>Escherichia coli</u>", J. Bacteriol., 172:p.6704-6712 (1990) and Japanese Patent Application laid-open No. HEI 3-58786 (Japanese Patent Application filing No. HEI 2-53255): "A DNA chain useful for synthesis of carotenoids" by the present inventors) having the carotenoid synthesis genes except for <u>crt</u>Z derived from <u>Erwinia uredovora</u>, is digested with <u>Bst</u>EII, treated with Klenow enzyme and religated to inactivate the <u>crt</u>X gene by frame shift. After that, the 6.0 kb <u>Asp</u>718(KpnI)-EcoRI fragment containing <u>crt</u>E, <u>crt</u>B, <u>crt</u>I and <u>crt</u>Y genes necessary for β-carotene production is taken out. The fragment is then inserted into the <u>Eco</u>RV sites of the <u>E. coli</u> vector pACYC184 and the desirable plasmid(named pACCAR16ΔcrtX, FIGURE 10) is obtained. <u>E. coli</u> containing this plasmid (pACCAR16ΔcrtX) is chloramphenicol resistant and has yellowish color due to β-carotene production.

Then, the plasmid pCAR16 is digested with <u>BstEll/Sna</u>Bl, treated with Klenow enzyme and religated to remove the 2.26 kb <u>BstEll-Sna</u>Bl fragment containing <u>crt</u>X and <u>crt</u>Y genes. After that, the 3.75 kb <u>Asp718(Kpnl)-Eco</u>Rl fragment containing <u>crt</u>E, <u>crt</u>B and <u>crt</u>I genes necessary for lycopene production is taken out. The fragment is then inserted into the <u>Eco</u>RV sites of the <u>E. coli</u> vector pACYC184 and the desirable plasmid(named pACCRT-EIB, FIGURE 10) is obtained. <u>E. coli</u> containing pACCRT-EIB is chloramphenicol resistant and has reddish color due to lycopene production (Cunningham Jr., F. X., Chamovitz, D., Misawa, N., Gatt, E., Hirschberf, J., "Cloning and functional expression in <u>Escherichia coli</u> of a cyanobacterial gene for lycopene cyclase, the enzyme that catalyzes the biosynthesis of β-carotene", FEBS Lett., 328: 130-138 (1993)).

Then, the plasmid pCAR16 is digested with <u>BstEII/Eco</u>52I, treated with Klenow enzyme and religated to remove the 3.7 kb <u>BstEII-Eco</u>52I fragment containing <u>crt</u>X, <u>crt</u>Y and <u>crt</u>I genes. After that, the 2.3 kb <u>Asp</u>718(<u>KpnI</u>)-<u>Eco</u>RI fragment containing <u>crt</u>E and <u>crt</u>B genes(FIGURE 2) necessary for phytoene production is taken out. The fragment is then inserted into the <u>Eco</u>RV sites of the <u>E. coli</u> vector pACYC184 and the decibel plasmid(named pACCRT-EB, FIGURE 10) is obtained. <u>E. coli</u> containing pACCRT-EB is chloramphenicol resistant and does not show color change as phytoene is colorless (Linden, H., Misawa, N., Chamovitz, D., Pecker, I., Hirschberg, J., Sandmann, G., "Functional complementation in <u>Escherichia coli</u> of different phytoene desaturase genes and analysis of accumulated carotenes", Z. Naturforsch. 46c: 1045-1051 (1991)).

(EXAMPLE 6) Screening of genes that increase β-carotene production

As the <u>E. coli</u> strain JM101 containing the above plasmid pACCAR16ΔcrtX shows yellowish color due to β-carotene production, it was investigated whether more yellowish transformant can be obtained by introducing cDNA expression library of <u>Phaffia rhodozyma</u> or <u>Haematococcus pluvialis</u>. As a first step, competent cells of <u>E. coli</u> JM101 containing pACCAR16ΔcrtX are prepared according to the method described in "Molecular cloning 2nd edition: Cold Spring Harbor Laboratory, 1.21-1.41(1989). Then, one hundred ng of each cDNA expression library of <u>Ph. rhodozyma</u> and <u>Ha. pluvialis</u> is introduced to 1 ml of the competent cells. Approx. 200,000 transformants of <u>Ph. rhodozyma</u> and approx. 40,000 transformants of <u>Ha. pluvialis</u> are obtained and inoculated for screening on the LB plate(bactotrypton 1%, yeast extract 0.5%, NaCl 1%, agar 1.5%) containing 150 μg/ml of ampicillin, 30 μg/ml of chloramphenicol and 1 mM of IPTG. From the screening, 5 strains of <u>Ph. rhodozyma</u> and 10 strains of <u>Ha. pluvialis</u> shows deep yellowish color than other strains and they are isolated. The plasmid DNA extracted from these strains is subject to restriction enzyme analysis, and it was found that the plasmids from the five strains and ten strains have common DNA fragment respectively. Of these screened plasmids derived from the cDNA expression libraries, a plasmid from <u>Ph. rhodozyma</u> was named pRH1(Figure 11) and another plasmid from <u>Ha. pluvialis</u> was named pHP1. In addition to that, a fragment is taken out after digesting pHP1 with <u>Sal</u>I and <u>Not</u>I, and then, the fragment is inserted into pBluescript KS+. The resulted plasmid was named pHP11(FIGURE 11) and was used for the experiments mentioned below.

(EXAMPLE 7) Nucleotide sequence determination on the gene that increases β-carotene production

From the plasmids pRH1 and pHP1, the deletion plasmids which lack various lengths from the original plasmids are prepared by the following procedures. By using those deletion plasmids, the nucleotide sequences are determined.

Decomposition of pRP1 is performed with both <u>Eco</u>RI and <u>Pst</u>I, or with both <u>Not</u>I and <u>Sph</u>I. Decomposition of pHP1 is performed with both <u>Aat</u>II and <u>Bam</u>HI, or with both <u>Kpn</u>I and <u>Eco</u>RI. After extraction with phenol/chloroform, DNA is recovered by ethanol precipitation. Each DNA fraction is then dissolved in 100 µI portions of ExoIII buffer(50mM Tris-HCI, 100mM NaCI, 5mM MgCl₂, 10mM 2-mercaptoethanol, pH 8.0) and is kept at 37°C after addition or 180 units of ExoIII nuclease. Ten µI portions of the solution are sampled every 30 seconds and transferred to tubes containing 10 µI of MB buffer(40 mM NaCI, 2 mM ZnCl₂, 10% glycerol, pH 4.5) in an ice bath. After sampling, the 10 tubes are kept at 65°C for 10 minutes to inactivate the enzyme. Then, 5 units of mung bean nuclease is added and kept at 37°C for 30 minutes. From one original plasmid, ten different kind of DNA fragments are recovered by agarose gel electrophoresis. The degree of deletion of each fragment varies. The terminals of the recovered DNAs are smoothed with Klenow enzyme to subject to ligation reaction at 16°C overnight, and by using resulting DNA, <u>E. coli</u> DH5α is transformed to obtain clones. The plasmids are prepared from the various clones obtained, and nucleotide sequences are determined by using luminescence primer cycle sequence kit(Applied Biosystems corp.) with an automatic sequencer.

As a result, it was found that the nucleotide sequence of the cDNA in pRH1 derived from Phaffia rhodozyma consists of 1,099 base pairs (SEQUENCE ID No. 4), and there is an open reading frame which encodes a polypeptide having 251 amino acids (which corresponds the region from A to B in Figures 4 and 5). It was also found that the nucleotide sequence of the cDNA in pHP1 derived from Haematococcus pluvialis consists of 1,074 base pairs (SEQUENCE ID No. 5), and there is an open reading frame which encodes a polypeptide having 259 amino acids (which corresponds the region from C to D in Figures 6 and 7). The amino acid sequences expected from these open reading frames are investigated by analyzing homology in the Gene Bank. Both of the amino acid sequences of Ph. rhodozyma and Ha. pluvialis have significant homology with the IPP isomerase gene of Saccharomyces cerevisiae, 27.0% for Ph. rhodozyma and 20.3% for Ha. pluvialis. Therefore the genes were identified as the IPP isomerase gene.

(EXAMPLE 8) Preparation of total DNA in Saccharomyces cerevisiae

Preparation of total DNA in <u>Saccharomyces cerevisiae</u> is performed according to the method described in "Methods in Yeast Genetics; a laboratory course manual: Cold Spring Harbor Laboratory, p.131-132(1990). <u>Sa. cerevisiae</u> S288C strain is inoculated in 10 ml of YPD media and cultured at 30°C overnight. The cultured cells are collected and suspended in 0.5 ml of sterilized water for washing. By discarding the supernatant, the yeast are collected again. A 0.2 ml of the mixture(2% Triton X-100, 1% SDS, 100 mM NaCl, 10 mM Tris-HCl(pH 8), 1mM EDTA), 0.2 ml of phenol/chloroform/isoamylalcohol (25/24/1 (v/v/v)) and 0.3 g of glass beads are added. After vortex mix for 3-4 minutes, two hundred μl of TE buffer(10 mM Tris-HCl(pH 8), 1 mM EDTA) is added. Then the solution is centrifuged for 5 minutes, and the supernatant is transferred to another tube and 1 ml of ethanol is added. Then the solution is centrifuged again for 2 minutes. The resulted pellet is dissolved in 0.4 ml of TE buffer. Then, two μl of RNase A(10 mg/ml) is added and the solution is stood for 5 minutes at 37°C. Then, ten μl of 4 M ammonium acetate and 1 ml of ethanol are added. After mixing well, the solution is centrifuged for 2 minutes and the resulted pellet is recovered. After drying the pellet, it was dissolved with 50 μl of TE buffer to have total DNA of <u>S. cerevisiae</u> S288C strain. By this preparation procedure, 3.4 μg of total DNA was obtained.

(EXAMPLE 9) Isolation of the IPP isomerase gene of Saccharomyces cerevisiae by PCR method

Based on the nucleotide sequence of the IPP isomerase gene of <u>S. cerevisiae</u> reported in the aforementioned reference(Anderson, M. S., Muehlbacher, M., Street, I.P., Profitt, J., Poulter, C. D., "Isopentenyl diphosphate: dimethylallyl diphosphate isomerase - an improved purification of the enzyme and isolation of the gene from <u>Saccharomyces cerevisiae</u>", J. Biol. Chem., 264:19169-19175(1989)), the primers below were synthesized.

Primer No. 1 5'-TCGATGGGGGTTGCCTTTCTTTTCGG-3'

Primer No. 2 5'-CGCGTTGTTATAGCATTCTATGAATTTGCC-3'

The procedure was designed to obtain PCR amplified IPP isomerase gene having <u>Tag</u>I sites at the upstream terminal and <u>Acc</u>II region at the downstream terminal. Thirty cycles of PCR is performed with 200 ng of total DNA of <u>S. cerevisiae</u> and PfuDNA polymerase (STRATAGENE). To express the IPP isomerase gene obtained by PCR in <u>E. coli</u>, it is digested with both <u>Tag</u>I and <u>Acc</u>II. Then, the gene was inserted into <u>Cla</u>I sites and <u>Sma</u>I sites of pBluescript KS+vector. The resulted plasmid was named pSI1(Figure 11). This DNA derived from <u>S. cerevisiae</u> had a nucleotide sequence consisting of 1,058 bp (SEQUENCE ID No. 6), and contained a gene which encodes IPP isomerase consisting of 288 amino acids(corresponds from E to F in Figures 8 and 9).

45

(EXAMPLE 10) Increase of lycopene production amount by introducing the IPP isomerase gene

Into the lycopene-producing <u>E. coli</u> JM101 strain (abbreviated as L hereafter) which contains pACCRT-EIB(Figure 10). pSPORT1 vector, pRH1 plasmid containing the IPP isomerase gene of <u>Phaffia rhodozyma</u>, pHP11 plasmid containing the IPP isomerase gene of <u>Haematococcus pluvialis</u> or pSI1 plasmid containing the IPP isomerase gene of <u>Saccharomyces cerevisiae</u>(FIGURE 11) are introduced respectively. These <u>E. coli</u> transformants are then plated on the LB plate containing 150 μg/ml of ampicillin(Ap), 30 μg/ml of chloramphenicol(Cm) and 1 mM of IPTG, and cultured at 28°C overnight. The three strains, in which each IPP isomerase gene were introduced, showed deep reddish color due to lycopene production compared with the control (lycopene-producing E.coli) in which only vector is introduced. Furthermore, growth rate of the three strains on agar plates were faster than the control strains and they always showed larger colonies than those of the control during culture. It is considered that due to introduction and expression of the IPP isomerase gene, the upstream of the biosynthetic pathway up to FPP became more efficient(see FIGURE 1), and consequently, increase of FPP supply led to increase of lycopene. As for faster growth rate, it is also considered that due to increase of FPP, sufficient amount of the substrate can be supplied not only for lycopene production but also for the production of other membrane components derived from FPP, that is, FPP or GGPP binding protein, and these components are necessary for growth of <u>E. coli</u>.

Increase of lycopene production amount by E.coli carrying the IPP isomerase gene is also confirmed by liquid culture. After overnight shaking culture of the LB media(5 ml) containing both Ap and Cm at 28°C, 2 ml of the media is taken and transferred to 200 ml of 2YT culture media(1.6% bactotrypton, 1% yeast extract, 0.5% NaCl) containing Ap, Cp and 0.1 mM IPTG, and shaking culture is performed at 230 rpm, 28°C. Five mI each of the media is sampled several hours' intervals to determine growth rate and lycopene content. Growth rate is calculated from absorbance at 650 nm. Lycopene content is determined according to the following procedure. The cells collected by centrifugation, 2.5 ml of acetone is added to the cells and stand for 30 minutes. Vortex mix once in a while. After filtration, absorbance at 474 nm is measured to determine the lycopene content based on the absorbance 185.0 for 1 mM lycopene (light path: 1 cm). JASCO UVIDEC-220B spectrophotometer is used. By using HPLC, it is confirmed that these strains actually produced lycopene and absorbance at 474 nm is attributable to lycopene. HPLC conditions are mentioned in EXAMPLE 11. The results are shown in Figure 12(growth curve) and Figure 13(lycopene production curve). As for the growth rate(Figure 12), there is no difference among any the strains including the control strains. This result is different from that obtained from culture plates. Probably, when the liquid culture is performed, even in the control strain which does not have exogenous IPP isomerase gene can grow rapidly, because the supply of the substrate for membrane components such as FPP and GGPP binding protein is enough compared to agar culture is done. In contrast, there is a big difference between the control strain having no exogenous IPP isomerase gene and the three exogenous IPP isomerase gene-carrying strains. During culture, the three strains always showed several times higher lycopene production amount compared with the control strain. Lycopene production amount per E. coli dry weight at 28 hr after the start of the culture is shown in Figure 14. The three strains containing the IPP isomerase gene showed 3.6-4.5 times higher production than the control strain. Lycopene-producing E. coli containing pHP11 is able to produce 1.03 mg lycopene per 1g dry weight.

(EXAMPLE 11) Increase of β-carotene production amount by introducing the IPP isomerase gene

Into the β -carotene producing \underline{E} . \underline{coli} JM101 strain (abbreviated as β hereafter) which contains pACCAR16 Δ crtX(FIGURE 10), either pSPORT1 vector or pRH1 plasmid containing the IPP isomerase gene of $\underline{Phaffia}$ $\underline{rhodozyma}$ is introduced separately. After overnight shaking culture of the LB media(5 ml) containing both Ap and Cm at 28°C, 1 ml of the media is taken and transferred to 100 ml of 2YT media containing Ap, Cm and 0.1 mM IPTG, and shaking culture is performed at 230 rpm at 28°C for 28 hr. The bacteria are collected by centrifugation and washed with 0.85% NaCl. After washing, the bacteria are suspended in 40 ml of acetone and allowed to stand for 30 minutes. Vortex mix once in a while. After filtration, absorbance at 454 nm is measured to determine β -carotene content based on the absorbance 134.4 for 1 mM β -carotene (light path: 1 cm). The result is shown in FIGURE 14. β -Carotene producing \underline{E} . \underline{coli} containing pRH1 produced 709 μ g of β -carotene per 1g dry weight. This amount is 1.5 times higher than the control.

By using HPLC on the above acetone extract, it is confirmed that these strains actually produced β -carotene and absorbance at 454 nm is attributable to β -carotene. Novapack HR 6μ C18(3.9 x 300 mm, Waters) is used as a column. Acetonitrile/methanol/2-propanol(90/6/4(v/v/v)) is used as an elution solvent. A photodiode array detector 996(Waters) is used to monitor an elution profile. The results showed that almost 100% of a peak appeared in a visible spectrum is β -carotene. As the β -carotene standard preparation, chemically synthesized β -carotene (Sigma) is used.

(EXAMPLE 12) Increase of phytoene production amount by introducing the IPP isomerase gene

Into the phytoene producing <u>E. coli</u> JM101 strain (abbreviated as P hereafter) which contains pACCRT-EB(FIGURE 10), any of pSPORT1 vector, pRH1 plasmid containing the IPP isomerase gene of <u>Phaffia rhodozyma</u> or pHP11 plas-

mid containing the IPP isomerase gene of <u>Haematococcus pluvialis</u> is introduced separately. After overnight shaking culture of the LB media(5 ml) containing both Ap and Cm at 28°C, 1 ml of the media is taken and transferred to 100 ml of 2YT media containing Ap, Cm and 0.1 mM IPTG, and shaking culture is performed at 230 rpm at 28°C for 28 hr. The bacteria are collected by centrifugation and washed with 0.85% NaCl. After washing, the bacteria are suspended in 40 ml of acetone and allowed to stand for 30 minutes. Vortex mix once in a while. After filtration and drying by rotary evaporator, partition is performed with 40 ml of petroleum ether and water. Absorbance of the ether layer at 286 nm is measured to determine phytoene content based on the absorbance 41.2 for 1 mM phytoene (light path: 1 cm). As HPLC analysis described in EXAMPLE 11 showed that 70% of the absorbance at 286 nm is attributable to phytoene, an and also actual phytoene content is adjusted to 70% of the above value. The result is shown in FIGURE 14. Phytoene-producing <u>E. coli</u> containing the IPP isomerase gene produced 1.7-2.1 times higher phytoene than control strain.

From the above examples, we showed that by introducing the IPP isomerase gene into β -carotene, lycopene or phytoene-producing <u>E. coli</u>, several times higher carotenoid production is actually achieved. It is considered that due to introduction and expression of the IPP isomerase gene, upstream of the biosynthetic pathway up to FPP became more efficient(see FIGURE 1), and consequently, increase of FPP supply led to increase of these carotenoids. Therefore, it is considered that this findings can be applicable not only for β -carotene, lycopene and phytoene productions but also for all other carotenoids such as astaxanthin and zeaxanthin.

The present invention provides a DNA chain which can significantly increase carotenoid production in biosynthesis of carotenoid by microorganisms and a method to obtain several times higher carotenoid production amount by introducing and expressing said DNA chain into carotenoid-producing microorganisms. It is expected that said DNA chain can be applicable to increase production in microorganisms not only for carotenoids but also for terpenoids and so forth which require same substrate(FPP) as carotenoids.

	[SEC	QUENC	CE L	ISTI	1G]											
	s	EQUE	NCE	ID N	o.:	1										
5	L	ENGT	н: 2	51												
	S	EQUE	NCE	TYPE	: am	ino	acid									
	T	OPOL	OGY:	lin	ear											
10	м	OLEC	ULAR	TYP	E: p	epti	de									
		EQUE			•	•										
		-		Pro	Asn	Ile	Val	Pro	Pro	Ala	Glu	Val	Arq	Thr	Glu	Gly
15					5					10			•		15	_
	Leu	Ser	Leu	Glu	Glu	Tyr	Asp	Glu	Glu	Gln	Val	Arg	Leu	Met	Glu	Glu
				20					25					30		
20	Arg	Cys	Ile 35	Leu	Val	Asn	Pro	Asp 40	Asp	Val	Ala	Tyr	Gly 45	Glu	Ala	Ser
	Lys	Lys	Thr	Cys	His	Leu	Met	Ser	Asp	Ile	Asn	Ala	Pro	Lys	Asp	Leu
		50					55					60				
25		His	Arg	Ala	Phe		Val	Phe	Leu	Phe		Pro	Ser	Asp	Gly	
	65 Leu	Leu	Leu	Gln	Arg	70 Arg	Ala	Asp	Glu	Lvs	75 Tle	Thr	Phe	Pro	Glv	80 Met
					85	••••			014	90				110	95	1100
30	Trp	Thr	Asn	Thr	Cys	Cys	Ser	His	Pro	Leu	Ser	Ile	Lys	Gly	Glu	Val
				100					105					110		
	Glu	Glu		Asn	Gln	Ile	Gly	Val	Arg	Arg	Ala	Ala	Ser	Arg	Lys	Leu
	G3	77.5.	115	T	G1	** - 7	D	120	G	0	D b	5	125		•	
35	GIU	130	GIU	rea	GIÀ	vaı	135	Thr	ser	Ser	Thr	140	PIO	Asp	ser	Pne
	Thr		Leu	Thr	Arg	Ile		Tyr	Leu	Ala	Pro		Asp	Gly	Leu	Trp
	145				-	150		-			155		•	•		160
40	Gly	Glu	His	Glu	Ile	Asp	Tyr	Ile	Leu	Phe	Ser	Thr	Thr	Pro	Thr	Glu
					165		_			170					175	
	His	Thr	Gly	180	Pro	Asn	Glu	Val	Ser 185	Asp	Thr	Arg	Tyr	Val 190	Thr	Lys
4 5	Pro	Glu	T.e.u		Δla	Mo+	Phe	Glu		Glu	Sor	Aen	Sar	Phe	Thr	Dro
		014	195	U 1	1114		1 1	200	ımp	0,1	DCI	11011	205	1110	1111	110
	Trp	Phe	Lys	Leu	Ile	Ala	Arg	Asp	Phe	Leu	Phe	Gly	Trp	Trp	Asp	Gln
50		210					215					220				
· -		Leu	Ala	Arg	Arg		Glu	Lys	Gly	Glu		Asp	Ala	Lys	Ser	
	225					230					235					240

Glu Asp Leu Ser Asp Asn Lys Val Trp Lys Met ***

5																
10		EQUE			0.:	2										
	L	ENGT	'H: 2	59												
	S	EQUE	NCE	TYPE	: am	ino	acid									
_	Т	OPOL	OGY:	lin	ear											
15	М	OLEC	ULAR	TYP	E: p	epti	de									
	s	EQUE	NCE:													
		Gln			Ala	Glu	Asp	Arg	Thr	Asp	His	Met	Arg	Gly	Ala	Sei
20					5					10					15	
	Thr	Trp	Ala	Gly	Gly	Gln	Ser	Gln	Asp	Glu	Leu	Met	Leu	Lys	Asp	Gl
				20					25					30		
25	Cys	Ile		Val	Asp	Ala	Asp	_	Asn	Ile	Thr	Gly		Val	Ser	Ly
	_		35	•	_		_	40	1		_		45			
	Leu	Glu 50	Cys	His	Lys	Phe	Leu 55	Pro	His	Gln	Pro	Ala 60	Gly	Leu	Leu	His
	Arq	Ala	Phe	Ser	Val	Phe		Phe	Asp	Asp	Gln		Arg	Leu	Leu	Lei
30	65					70			•	•	75	•	,			8
	Gln	Gln	Arg	Ala	Arg	Ser	Lys	Ile	Thr	Phe	Pro	Ser	Val	Trp	Thr	Ası
					85					90					95	
35	Thr	Cys	Cys		His	Pro	Leu	His	_	Gln	Thr	Pro	Asp		Val	Asp
				100					105					110		
	Gln	Leu		Gln	Val	Ala	Asp	_	Thr	Val	Pro	Gly		Lys	Ala	Ala
	37.0	T1-	115	T -	.	61	•••	120	-	63	-1-		125		63	_
40	Ald	11e	Arg	rys	геп	GIU	135	GIU	ьeи	GIÀ	TTE	140	ALA	HIS	GIN	тел
		100					100					140				

Pro Ala Ser Ala Phe Arg Phe Leu Thr Arg Leu His Tyr Cys Ala Ala

Asp Val Gln Pro Ala Ala Thr Gln Ser Ala Leu Trp Gly Glu His Glu

Met Asp Tyr Ile Leu Phe Ile Arg Ala Asn Val Thr Leu Ala Pro Asn

Pro Asp Glu Val Asp Glu Val Arg Tyr Val Thr Gln Glu Glu Leu Arg

		210					215			Glu		220				_
5	Ile	Ile	Ala	Ala	Arg	Phe	Leu	Glu	Arg	Trp	Trp	Ala	Asp	Leu	Asp	Ala
	225					230					235					240
	Ala	Leu	Asn	Thr	Asp	Lys	His	Glu	Asp	Trp	Gly	Thr	Val	His	His	Ile
					245					250					255	
10	Asn	Glu	Ala	***												
15	S	EQUE	NCE	ID N	io.:	3										
	L	ENGT	'H: 2	88												
	S	EQUE	NCE	TYPE	: am	uno	acid	l								
20	T	OPOL	OGY:	lin	ear											
	M	OT EC	111 A E				٠.									
	m	OLEC	ULAR	CTIP	E: p	epti	.ae									
25		EQUE														
23	Met	Thr	Ala	Asp			Ser	Met	Pro	His	Gly	Ala	Val	Ser	Ser	Tyr
		_	_	-	5					10					15	
	Ala	Lys	Leu		Gln	Asn	Gln	Thr		Glu	Asp	Ile	Leu		Glu	Phe
30	D==	C1	71.	20	D	•	63 -	~ 1	25	_	_			30	_	_
	PIO	GIU	35	me	PIO	rea	GIN		Arg	Pro	Asn	Thr	-	Ser	Ser	Glu
	Thr	Sor) cn	Glu	505	C111	40	mb~	Cys	Dho	C	45	***		63
	1111	50	ASII	ASP	GIU	ser	55 55	Glu	THE	cys	Pne	Ser 60	GTÀ	HIS	Asp	GIU
35	Glu		Tle	Lvs	T.eu	Mot		Glu	Δen	Cys	Tla		LOU	λcn	ш-г-г	N C D
	65			_,_	200	70	71011	014	AD II	Cys	75	Vai	Ten	vab	пр	AS P
	Asp	Asn	Ala	Ile	Gly		Glv	Thr	Lvs	Lys		Cvs	His	Leu	Met	
40	•				85		•		_	90		- 2 -			95	
	Asn	Ile	Glu	Lys	Gly	Leu	Leu	His	Arg	Ala	Phe	Ser	Val	Phe		Phe
				100										110		
	Asn	Glu	Gln	Gly	Glu	Leu	Leu	Leu	Gln	Gln	Arg	Ala	Thr	Glu	Lys	Ile
45			115					120					125		_	
	Thr	Phe	Pro	Asp	Leu	Trp	Thr	Asn	Thr	Cys	Cys	Ser	His	Pro	Leu	Cys
		130					135					140				
	Ile	Asp	Asp	Glu	Leu	Gly	Leu	Lys	Gly	Lys	Leu	Asp	Asp	Lys	Ile	Lys
50	145					150					155					160
	Gly	Ala	Ile	Thr	Ala	Ala	Val	Arg	Lys	Leu	Asp	His	Glu	Leu	Gly	Ile

					165					176					175		
	Pro	Glu	Asp	Glu	Thr	Lys	Thr	Arg	Gly	Lys	Phe	His	Phe	Leu	Asn	Arg	
5				180					185					190			
	Ile	His		Met	Ala	Pro	Ser		Glu	Pro	Trp	Gly		His	Glu	Ile	
) en	Tyr	195	T.e.u	Dha	ጥላታ	Luc	200	Aen	λla	Lve	Glu	205 Asn	Len	Thr	Val	
10	vəħ	210	116	Deu	1110	- 7 -	215	116	no II	ALG	D _I 3	220	non	Deu	1111	Vai	
	Asn	Pro	Asn	Val	Asn	Glu	Val	Arg	Asp	Phe	Lys	Trp	Val	Ser	Pro	Asn	
	225					230					235					240	
15	Asp	Leu	Lys	Thr		Phe	Ala	Asp	Pro		Tyr	Lys	Phe	Thr	Pro	Trp	
					245					250		_	_		255		
	Phe	Lys	Ile	11e 260	Cys	Glu	Asn	Tyr	265	Phe	Asn	Trp	Trp	Glu 270	Gln	Leu	
	ASD	Asp	Len		Glu	Val	Glu	Asn		Ara	Gln	Tle	His		Met	Leu	
20			275	552	014			280		,	J		285	,		201	

25																	
	ς	EQUE	NCF	מ חז	o •	4											
					•••	•											
30		ENGT															
	S	EQUE	NCE	TYPE	: nu	clei	c ac	id									
	S	TRAN	DNES	S: d	oubl	е											
35	Т	OPOL	OGY:	lin	ear												
	М	OLEC	ULAR	TYP	E: c	DNA											
	0	RIGI	N														
40		ORC	ANIS	M: I	hafi	ia n	hodo	zyma	l								
40		STI	RAIN:	ATO	C 24	230											
	S	EQUE					TC										
	J					CODE		ne.									
4 5							CL	,5									
		LOC	CATI	.ON:	99	821											
		DET	rerm]	NATI	ON M	IETH(D: E	E									
50		EQUE															
	CCC	ACGC	GTC (CGCA	CATC	rc Go	CATAT	TATCA	CTI	TCCI	rcct	TCC	AGAA	CAA (STTCI	GAGTC	60

	AAC	CGAA	AAG Z	AAAG	AAGG	CA G	AGGA	AAAT	A TA	TTCT.	AG A	rg T	CC A	rg c	CC A	AC AT	т 116
5												Met :	Ser 1	let I	Pro A	sn Il	е
	Cutan	ccc	ccc	ccc	CNC	CTC	CCN	3.00	C	CCN	cmc	AGT	~~	<i>~</i> • • • • • • • • • • • • • • • • • • •	63.6	5	1.64
																	164
	val	PIO	PIO		GIU	vaı	AIG	Thr			Leu	Ser	Leu		GIU	Tyr	
10	C > T	616	C. C.	10	000		050		15					20			
												TGT					212
	Asp	GIU		GIN	vai	Arg	Leu		Glu	Glu	Arg	Cys		Leu	Val	Asn	
	222	212	25					30					35				
15												AAG					260
	Pro		Asp	Val	Ala	Tyr		Glu	Ala	Ser	Lys	Lys	Thr	Cys	His	Leu	
		40					45					50					
												CAC					308
20		Ser	Asn	Ile	Asn		Pro	Lys	Asp	Leu	Leu	His	Arg	Ala	Phe	Ser	
	55					60					65					70	
												CTG					356
	Val	Phe	Leu	Phe	Arg	Pro	Ser	Asp	Gly	Ala	Leu	Leu	Leu	Gln	Arg	Arg	
25					75					80					85		
												ACC					404
	Ala	Asp	Glu	Lys	Ile	Thr	Phe	Pro	Gly	Met	Trp	Thr	Asn	Thr	Cys	Cys	
				90					95					100			
30												GAG					452
	Ser	His	Pro	Leu	Ser	Ile	Lys	Gly	Glu	Val	Glu	Glu	Glu	Asn	Gln	Ile	
			105					110					115				
0.5	GGT	GTT	CGA	CGA	GCT	GCG	TCC	CGA	AAG	TTG	GAG	CAC	GAG	CTT	GGC	GTG	500
35	Gly	Val	Arg	Arg	Ala	Ala	Ser	Arg	Lys	Leu	Glu	His	Glu	Leu	Gly	Val	
		120					125					130					
	CCT	ACA	TCG	TCG	ACT	CCG	CCC	GAC	TCG	TTC	ACC	TAC	CTC	ACT	AGG	ATA	548
40	Pro	Thr	Ser	Ser	Thr	Pro	Pro	Asp	Ser	Phe	Thr	Tyr	Leu	Thr	Arg	Ile	
70	135					140					145					150	
	CAT	TAC	CTC	GCT	CCG	AGT	GAC	GGA	CTC	TGG	GGA	GAA	CAC	GAG	ATC	GAC	596
	His	Tyr	Leu	Ala	Pro	Ser	Asp	Gly	Leu	Trp	Gly	Glu	His	Glu	Ile	Asp	
45					155					160					165		
	TAC	ATT	CTC	TTC	TCA	ACC	ACA	CCT	ACA	GAA	CAC	ACT	GGA	AAC	CCT	AAC	644
	Tyr	Ile	Leu	Phe	Ser	Thr	Thr	Pro	Thr	Glu	His	Thr	Gly	Asn	Pro	Asn	
				170					175					180			
50	GAA	GTC	TCT	GAC	ACT	CGA	TAT	GTC	ACC	AAG	ccc	GAG	CTC	CAG	GCG	ATG	692
												Glu					

	185	190	195	
	TTT GAG GAC GAG TCT	AAC TCA TTT ACC	CCT TGG TTC AAG TTG	ATT GCC 740
5	Phe Glu Asp Glu Ser	Asn Ser Phe Thr	Pro Trp Phe Lys Leu	Ile Ala
	200	205	210	
			CAA CTT CTC GCC AGA	
10	215	Gly Trp Trp Asp	Gln Leu Leu Ala Arg 225	-
	7.7	77	TTG GAG GAT CTC TCG	230 GAC AAC 836
			Leu Glu Asp Leu Ser	
	235		240	245
15	AAA GTC TGG AAG ATG	TAGTCGACC CTTCT	TTCTG TACAGTCATC TCA	GTTCGCC 890
	Lys Val Trp Lys Met	***		
	250			
20	TGTTGGTTGC TTGCTTCTT	IG CTCTTCTTTC TA	TATATCTT TTTTCTTGCC	TGGGTAGACT 950
	ጥርልጥርጥጥርጥ አርአጥአርርአባ	דא רכראידאראידא רא	TAAACTCT ATTTCTTGTT	CTTT TO
	TONICITIES ACAINGCA	IA COCAIACAIA CA	IAAACICI AIIICIIGII	CITIAICICI 1010
25	CTTCTAAGGG AATCTTCAA	AG ATCAATTTCT TT	TTGGGCTA CAACATTTCA	GATCAATGTT 1070
	GCTTTTCAGA CTACAAAA	AA AAAAAAAA 10	99	
30				
35	SEQUENCE ID No.:	5		
	LENGTH: 1074			
	SEQUENCE TYPE: nu	cleic acid		
40	STRANDNESS: double	e		
	TOPOLOGY: linear			
	MOLECULAR TYPE: C	DNA		
45	ORIGIN			
	ORGANISM: Haema	tococcus pluvia	lis	
	STRAIN: NIES-14	44		
50	SEQUENCE CHARACTE	RISTIC		

17

CHARACTERISTIC CODE: CDS

	LOC	ATII	ON:	145.	.921												
	DET	ERMI	TAN	ON M	ETHO	D: E	Ξ										
S	EQUE	NCE:															
ATC	CTAC	CTT C	GAAC	CTG	sc co	GGC	GCA	TC	CGAT(GACG	CGA'	rgct	TCG	TTCG:	rtgc	TC	60
AGAC	GCC?	CA C	CGCAT	rttcc	CC CC	GCG:	rgaac	TC	CGCG	CAGC	AGC	CCAG	CTG	TGCA	CACG	CG	120
CGAC	יזיררו	ነርጥ ባ	የተልርረ	בררר:	G AZ	رور <u>،</u>	ነጥር (`AG (ጉጥር () بل مانات	3CC (SAG	CAC	CGC i	404	GAC	179
COIL		.01			.0 .1									Arg			1,,
											5		-			n	
CAT	ATG	AGG	GGT	GCA	AGT	ACC	TGG	GCA	GGC	GGG	CAG	TCG	CAG	GAT	GAG	22	2
His	Met	Arg	Gly	Ala	Ser	Thr	Trp	Ala	Gly	Gly	Gln	Ser	Gln	Asp	Glu		
				15					20					25			
														AAC			0
ren	met	Leu	TÀ2	Asp	GIU	Cys	TTE	Leu 35	Val	Asp	Ala	Asp	Asp 40	Asn	TTE		
ACA	GGC	CAT		AGC	AAG	CTG	GAG		CAC	AAG	TTC	СТА		CAT	CAG	31	8
														His			
		45					50					55					
CCT	GCA	GGC	CTG	CTG	CAC	CGG	GCC	TTC	TCT	GTA	TTC	CTG	TTT	GAC	GAC	36	6
Pro		Gly	Leu	Leu	His		Ala	Phe	Ser	Val		Leu	Phe	Asp	Asp		
a. a	60		ama	ama	ama	65	a. c	aam	60 3	<i>aa</i> .	70				mma	4.1	
														ACA Thr			4
75	Gry	Arg	Dea	Dea	80	01	0111	n g	ALG	85	501	Dys	110		90		
CCC	AGT	GTG	TGG	ACC	AAC	ACC	TGC	TGC	AGC	CAC	CCT	СТА	CAT	GGG	CAG	46	2
Pro	Ser	Val	Trp	Thr	Asn	Thr	Cys	Cys	Ser	His	Pro	Leu	His	Gly	Gln		
				95					100					105			
														ACA			0
Thr	Pro	Asp		Val	Asp	Gln	Leu		Gln	Val	Ala	Asp		Thr	Val		
CCT	ccc	CCN	110	CCM	CCM	ccc	N TO C	115	N N C	mmc	CAC	CAC	120	CMC	ccc	E E	0
														CTG Leu			0
110	01 1	125	273	.114	174 U		130	9	2,3	D Cu	724	135		204	J_1		
ATA	CCA		CAC	CAG	CTG	CCG		AGC	GCG	TTT	CGC			ACG	CGT	60	6
Ile	Pro	Ala	His	Gln	Leu	Pro	Ala	Ser	Ala	Phe	Arg	Phe	Leu	Thr	Arg		
	140					145					150						

	TTG	CAC	TAC	TGC	GCC	GCG	GAC	GTG	CAG	CCG	GCT	GCG	ACA	CAA	TCA	GCA	654
	Leu	His	Tyr	Cys	Ala	Ala	Asp	Val	Gln	Pro	Ala	Ala	Thr	Gln	Ser	Ala	
5	155					160					165					170	
		TGG															702
	Leu	Trp	Gly	Glu	His	Glu	Met	Asp	Tyr	Ile	Leu	Phe	Ile	Arg	Ala	Asn	
					175					180					185		
10		ACC															750
	Val	Thr	Leu		Pro	Asn	Pro	Asp		Val	Asp	Glu	Val		Tyr	Val	
				190		000	a	> mc	195	63.6	000	C1.C	3 3 m	200	mmc	CNN	700
15		CAG															798
	THI	Gln	205	GIU	ьеи	Arg	GIII	210	met	GIII	FIO	ASP	215	GIY	Den	GIII	
	ጥርር	TCG		ጥርር	ידיידי	CGC	ATC		GCC	GCG	CGC	TTC		GAG	CGC	TGG	846
		Ser															
20	•	220		•			225				-	230			_	_	
	TGG	GCT	GAC	CTA	GAC	GCG	GCC	CTG	AAC	ACT	GAC	AAA	CAC	GAG	GAT	TGG	894
	Trp	Ala	Asp	Leu	Asp	Ala	Ala	Leu	Asn	Thr	Asp	Lys	His	Glu	Asp	Trp	
25	235					240					245					250	
	GGA	ACG	GTG	CAT	CAC	ATC	AAC	GAA	GCG	TGA	AAA	CAG	AAGC	IGTA	GG 9	940	
	Gly	Thr	Val	His		Ile	Asn	Glu	Ala	***							
					255				3 N.M.	amma (COOT!	acms.	nom (- m	non a consc	1000
30	ATG	TCAA	JAC 1	ACGT	ATG	1G G(. عى ى د	1.1.00	_ AI	21164	3000	CII	JGIM.	ICI (-111.	ITACIO	1000
	AGA	CTGA	ACC '	TGCAG	CTG	SA GA	ACAAS	rggto	G AG	CCA	ATTC	AAC'	TTTC	CGC 1	rgcao	TGGA	1060
		01011															
35	AAA	AAAA	AAA .	AAAA	10	7 4											
40	5	EQUE	NCE	ID N	0.:	6											
	I	ENGT	H: 1	058													
	9	EOUE	NCE	TYPE	: nu	clei	c ac	id									
45	9	TRAN	IDNES	s: d	oubl	e											
		OPOL															
		OLEC				en∩m	ic D	NA									
50		RIGI			y	C11011											
		VVT GT															

		ORC	SANIS	5M: 5	Saccl	haron	nyces	s ce	revi	siae								
		STE	RAIN	: S21	88C													
5	S	EQUE	NCE	CHAR	ACTE	RIST	IC											
		CH	ARACI	reri:	STIC	CODE	E: CI	DS										
		LOC	CATI	ION:	187	105	50											
10		DET	rerm:	INAT:	ION I	METHO		S										
	s		NCE:															
	TCG	ATGG	GGG 1	TTGC	CTTT	CT T	TTTC	GGTC	T TA	ACTC	CATT	TAT	'ATTI	TTA	TATT	CATI	TT	60
15																		
	YTAT	CTAT	TTA A	ACAG	GAAA	CA G	TTTT	CTAG	T GA	CAAG	AAGG	CGI	TATA'	CCC	ACTT.	AATI	CA	120
	ATA	TTAG	AGT .	ATTC	GTAT	TT G	GAAT.	ACAG	G AA	GAGT	AAAA	ATA	AGCC	AAA	AATT	CATT	CAC	180
20																		
	ACC	rca i	ATG .	ACT	GCC	GAC 2	AAC .	AAT	AGT	ATG	CCC	CAT	GGT	GCA	GTA	TCT	AGT	23
			Met	Thr	Ala	Asp	Asn	Asn	Ser	Met	Pro	His	Gly	Ala	Val	Ser	Ser	
25							5					10					15	
											GAA							79
	Tyr	Ala	Lys	Leu			Asn	GIn	Thr		Glu	Asp) Ile	Leu			1	
	ىلىنلىش	רכתי	GAA	ע ידיים	20 בידים		מידים	CAA	CAA	25 AGA	CCT	አልጥ	י ארר	CCA	30 TCT		, ,	27
30											Pro							۲,
				35					40					45		001		
	GAG	ACG	TCA	AAT	GAC	GAA	AGC	GGA	GAA	ACA	TGT	TTT	TCI	GGT	CAT	GAT	3	75
35	Glu	Thr	Ser	Asn	Asp	Glu	Ser	Gly	Glu	Thr	Cys	Phe	Ser	Gly	His	Asp	•	
			50					55					60)				
	GAG	GAG	CAA	ATT	AAG	TTA	ATG	AAT	GAA	AAT	TGT	ATT	GTI	TTG	GAT	TGG	5 4	23
	Glu	Glu	Gln	Ile	Lys	Leu	Met	Asn	Glu	Asn	Cys	Ile	val	Leu	Asp	Trp)	
40		65					70					75						
											AAA							71
		Asp	Asn	Ala	Ile		Ala	Gly	Thr	Lys	Lys	Val	Cys	His	Leu			
45	80		3 000	<i>~</i>		85 CCT	777.3	Cm >	C > T	o c m	90		. maa	cma		95		
											GCA							19
	GIU	ASII	TIE	GIU	100	GIY	Leu	rea	nıs		Ala	Pne	ser	vai	110			
	ጥጥር	<u>አል</u> ጥ	GAA	CDD		GAA	ፈጥጥ እ	ىلغلى	מידיים	105 440	CAA	AC)	GC C	ልርጥ			ς,	67
50											Gln							· ,
				115	1	-14			120			9		125		_, .	-	
									•									

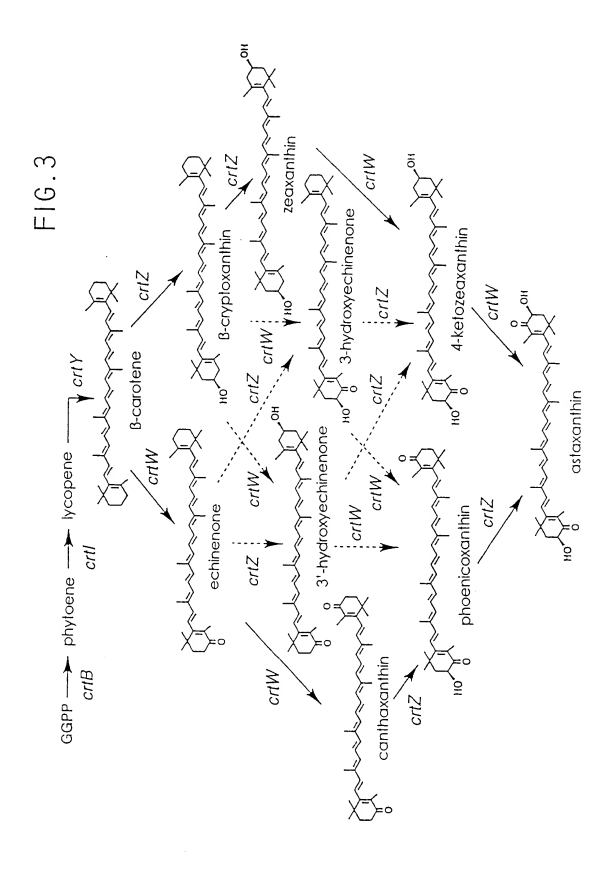
	ATA	ACT	TTC	CCT	GAT	CTT	TGG	ACT	AAC	ACA	TGC	TGC	1'CT	CAT	CCA	CTA	615
	Ile	Thr	Phe	Pro	Asp	Leu	Trp	Thr	Asn	Thr	Cys	Cys	Ser	His	Pro	Leu	
5			130					135					140				
	TGT	ATT	GAT	GAC	GAA	TTA	GGT	TTG	AAG	GGT	AAG	CTA	GAC	GAT	AAG	ATT	663
	Cys	Ile	Asp	Asp	Glu	Leu	Gly	Leu	Lys	Gly	Lys	Leu	Asp	Asp	Lys	Ile	
		145					150					155					
10	AAG	GGC	GCT	ATT	ACT	GCG	GCG	GTG	AGA	AAA	CTA	GAT	CAT	GAA	TTA	GGT	711
	Lys	Gly	Ala	Ile	Thr	Ala	Ala	Val	Arg	Lys	Leu	Asp	His	Glu	Leu	Gly	
	160					165					170					175	
	ATT	CCA	GAA	GAT	GAA	ACT	AAG	ACA	AGG	GGT	AAG	TTT	CAC	TTT	TTA	AAC	759
15	Ile	Pro	Glu	Asp	Glu	Thr	Lys	Thr	Arg	Gly	Lys	Phe	His	Phe	Leu	Asn	
					180					185					190		
	AGA	ATC	CAT	TAC	ATG	GCA	CCA	AGC	AAT	GAA	CCA	TGG	GGT	GAA	CAT	GAA	807
	Arg	Ile	His	Tyr	Met	Ala	Pro	Ser	Asn	Glu	Pro	Trp	Gly	Glu	His	Glu	
20				195					200					205			
	ATT	GAT	TAC	ATC	CTA	TTT	TAT	AAG	ATC	AAC	GCT	AAA	GAA	AAC	TTG	ACT	855
	Ile	Asp	Tyr	Ile	Leu	Phe	Tyr	Lys	Ile	Asn	Ala	Lys	Glu	Asn	Leu	Thr	
25			210					215					220				
	GTC	AAC	CCA	AAC	GTC	AAT	GAA	GTT	AGA	GAC	TTC	AAA	TGG	GTT	TCA	CCA	903
	Val	Asn	Pro	Asn	Val	Asn	Glu	Val	Arg	Asp	Phe	Lys	Trp	Val	Ser	Pro	
		225					230					235					
30	AAT	GAT	TTG	AAA	ACT	ATG	TTT	GCT	GAC	CCA	AGT	TAC	AAG	TTT	ACG	CCT	951
	Asn	Asp	Leu	Lys	Thr	Met	Phe	Ala	Asp	Pro	Ser	Tyr	Lys	Phe	Thr		
	240					245					250					255	
							GAG										999
35	Trp	Phe	Lys	Ile		Cys	Glu	Asn	Tyr		Phe	Asn	Trp	Trp		Gln	
					260					265					270		
																ATG 1	047
	Leu	Asp	Asp		Ser	Glu	Val	Glu		Asp	Arg	Gln	Ile		Arg	Met	
40				275					280					285			
		AAT	CAA	CG 1	058												
	Leu	***															

Claims

- 1. A DNA chain having characteristic of increasing carotenoid production, and containing the nucleotide sequence which encodes the polypeptide having the amino acid sequence substantially described in SEQUENCE ID No. 1, or a DNA chain which hybridizes with said DNA chain.
- 2. A DNA chain having characteristic of increasing carotenoid production, and containing the nucleotide sequence which encodes the polypeptide having the amino aid sequence substantially described in SEQUENCE ID No. 2, or a DNA chain which hybridizes with said DNA chain.
 - 3. A method for producing carotenoid characterized by introducing DNA chain described in one of claim 1 or 2 into carotenoid-producing microorganisms, culturing said transformed microorganism and obtaining higher carotenoid

content in the culture broth and cells .

4. A method for producing carotenoid characterized by introducing DNA chain containing the nucleotide sequence which encodes the polypeptide having the amino acid sequence substantially described in SEQUENCE ID No. 3, or DNA chain which hybridizes with said DNA chain introducing to carotenoid-producing microorganism, culturing said transformed microorganism and obtainig higher carotenoid content in the culture broth and cells.



```
18
                                  27
                                              36
                                                         45
 ATG TOO ATG COO AAC ATT GTT COO COO GCC GAG GTC CGA ACC GAA GGA CTC AGT
 Met Ser Met Pro Asn Ile Val Pro Pro Ala Glu Val Arg Thr Glu Gly Leu Ser
                      72
                                  81
 ITA GAA GAG TAC GAT GAG GAG CAG GTC AGG CTG ATG GAG GAG CGA TGT ATT CTT
 Leu Glu Glu Tyr Asp Glu Glu Gln Val Arg Leu Met Glu Glu Arg Cys Ile Leu
                     126
                                 135
                                             144
                                                        153
                                                                     162
 GTT AAC CCG GAC GAT GTG GCC TAT GGA GAG GCT TCG AAA AAG ACC TGC CAC TTG
Val Asn Pro Asp Asp Val Ala Tyr Gly Glu Ala Ser Lys Lys Thr Cys His Leu
         171
                     180
                                 189
                                             198
                                                        207
                                                                     216
ATG TCC AAC ATC AAC GCG CCC AAG GAC CTC CTC CAC CGA GCA TTC TCC GTG TTT
Met Ser Asn Ile Asn Ala Pro Lys Asp Leu Leu His Arg Ala Phe Ser Val Phe
                                 243
                                             252
                                                        261
CTC TTC CGC CCA TCG GAC GGA GCA CTC CTG CTT CAG CGA AGA GCG GAC GAG AAG
Leu Phe Arg Pro Ser Asp Gly Ala Leu Leu Leu Gln Arg Arg Ala Asp Glu Lys
                                                                     90
                    288
                                 297
                                            306
                                                                    324
ATT ACG TIC CCT GGA ATG TGG ACC AAC ACG TGT TGC AGT CAT CCT TTG AGC ATC
Ile Thr Phe Pro Gly Met Trp Thr Asn Thr Cys Cys Ser His Pro Leu Ser Il:
                    342
                                 351
                                                        369
ANG GGC GAG GTT GAA GAG GAG AAC CAG ATC GGT GTT CGA CGA GCT GCG TCC CGA
Lys Gly Glu Val Glu Glu Glu Asn Gln Ile Gly Val Arg Arg Ala Ala Ser Arg
                    396
                               405
                                            414
                                                                    432
AAG TTG GAG CAC GAG CTT GGC GTG CCT ACA TCG TCG ACT CCG CCC GAC TCG TTC
Lys Leu Glu His Glu Leu Gly Val Pro Thr Ser Ser Thr Pro Pro Asp Ser Phe
                    450
                                 459
                                            468
                                                       477
ACC TAC CTC ACT AGG ATA CAT TAC CTC GCT CCG AGT GAC GGA CTC TGG GGA GAA
Thr Tyr Leu Thr Arg Ile His Tyr Leu Ala Pro Ser Asp Gly Leu Trp Gly Glu
                    504
                                            522
CAC GAG ATC GAC TAC ATT CTC TTC TCA ACC ACA CCT ACA GAA CAC ACT GGA AAC
His Glu Ile Asp Tyr Ile Leu Phe Ser Thr Thr Pro Thr Glu His Thr Gly Asn
                                                                    180
                                567
                                            576
CCT AAC GAA GTC TCT GAC ACT CGA TAT GTC ACC AAG CCC GAG CTC CAG GCG ATG
Pro Asn Glu Val Ser Asp Thr Arg Tyr Val Thr Lys Pro Glu Leu Gln Ala Met
```

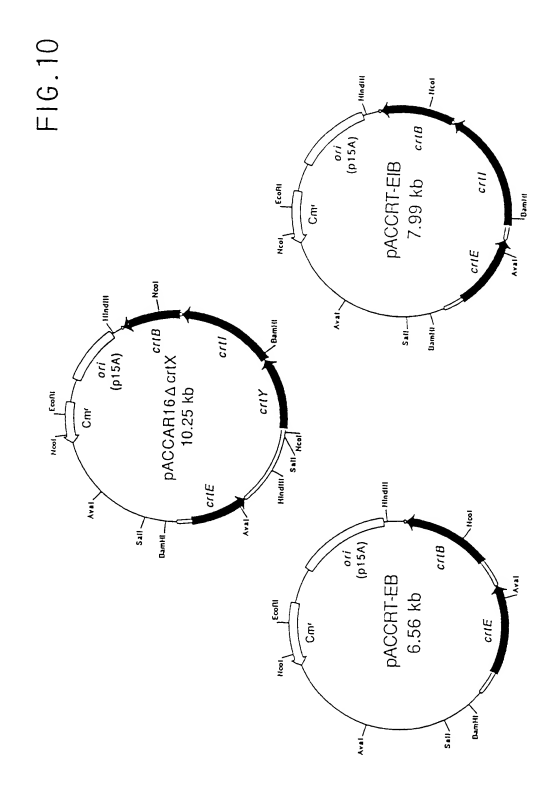
```
612 621 630 639
      603
TIT GAG GAC GAG TCT AAC TCA TIT ACC CCT TGG TTC AAG TTG ATT GCC CGA GAC
Phe Glu Asp Glu Ser Asn Ser Phe Thr Pro Trp Phe Lys Leu Ile Ala Arg Asp
                                                    216
              666
                                                    702
                      675
                                 684
                                          693
      657
TTC CTG TTT GGC TGG GAT CAA CTT CTC GCC AGA CGA AAT GAA AAG GGT GAG
Phe Leu Phe Gly Trp Trp Asp Gln Leu Leu Ala Arg Arg Asn Glu Lys Gly Glu
                       729
                                  738
               720
GTC GAT GCC AAA TCG TTG GAG GAT CTC TCG GAC AAC AAA GTC TGG AAG ATG TAG
В
```

С																	
Ĭ		9			18			27			36			45			54
ATG	CAG	-	CTT	GCC		GAC	CGC		GAC	CAT		λGG	GGT	_	AGT	ACC	-
Met	Gln	Leu	Leu	Ala	Glu	Asp	λrg	Thr	λзр	His	Met	Arg	Gly	Ala	Ser	Thr	Trp
																	18
		63			72			81			90			99			108
GCλ	GGC	GGG	CAG	TCG	ÇλG	GAT	GAG	CTG	ATG	CTG	λλG	GAC	GAG	TGC	ATC	TTG	GTG
Ala	GŢĀ	Gly	Gln	Ser	Gln	λsp	Glu	Leu	Met	Leu	Lys	dey	Glu	Cys	Ile	Leu	
																	36
		117	~ . ~		126		~~~	135			144		c	153	c		162
	GCT Ala																
АЗР	VT4	vab	хэр	V 311	116	1111	GIY	UT3	ATT	361	Lys	red	GIU	Cys	113	Lys	54
		171			180			189			198			207			216
CTA	CCA		CAG	CCT		GGC	CTG		CAC	CGG		TTC	TCT	GTA	TTC	CTG	
	Pro																
						_											72
		225			234			243			252			261			270
GAC	GAC	CAG	GGG	CGA	CTG	CIG	CTG	CAA	CAG	CGT	GCA	CGA	TCA	XXX	ATC	ACA	TTC
Asp	Αsp	Gln	Gly	λrg	Leu	Leu	Leu	Gln	Gln	λrg	Ala	Arg	Ser	Lys	Ile	Thr	
																	90
		279			288			297			306			315			324
	AGT																
PIO	Ser	VAI	Trp	ine	ASI	Int	Cys	CA2	Ser	uls	PLO	reu	ura	GLY	GIII	1111	108
		333			342			351			360			369			378
GAT	GAG		GAC	CAA		AGC	CAG		GCC	GAC		ACA	GTA		GGC	GCA	
	Glu																
										•	-				-		126
		387			396			405			414			423			432
GCT	GCT	GCC	ATC	CGC	λAG	TTG	GAG	CAC	GAG	CTG	GGG	ATA	CCA	GCG	CAC	CAG	CTG
Ala	λla	Ala	Ile	Arg	Lys	Leu	Glu	His	Glu	Leu	Gly	Ile	Pro	Ala	His	Gln	Leu
																	144
		441			450			459			468			477			486
	GCC																
Pro	Ala	Ser	Ala	Phe	yrg	Phe	Leu	Thr	λrg	Leu	HIS	Tyr	Cys	YIZ	VIA	qeA	
		*05			504			513			522			531			162 540
CNC	CCG	495				~C.	CCA		TCC	ccc		CAC	GAA		GAC	TAC	
	Pro																
GIM	.10	Ala	ALA	1111	GIII	361		Deu	1.p	u.,	014					-] -	180
		549			558			567			576			585			594
TTA	TTC		CGG	GCC		GTC	ACC		GCG	ccc		CCT	GAC	GAG	GTG	GAC	
	Phe																
			•														198

```
612 621 630 639 648
GTC AGG TAC GTG ACG CAG GAG GAG CTG CGG CAG ATG ATG CAG CCG GAC AAT GGG
Val Arg Tyr Val Thr Gln Glu Glu Leu Arg Gln Met Met Gln Pro Asp Asn Gly
                          675
                                     684
                666
TTG CAA TGG TCG CCG TGG TTT CGC ATC ATC GCC GCG CGC TTC CTT GAG CGC TGG
Leu Gln Trp Ser Pro Trp Phe Arg Ile Ile Ala Ala Arg Phe Leu Glu Arg Trp
                                   738 747
                           729
                720
TGG GCT GAC CTA GAC GCG GCC CTG AAC ACT GAC AAA CAC GAG GAT TGG GGA ACG
Trp Ala Asp Leu Asp Ala Ala Leu Asn Thr Asp Lys His Glu Asp Trp Gly Thr
                        780
       765
                 774
GTG CAT CAC ATC AAC GAA GCG TGA
Val His His Ile Asn Glu Ala;***
259
```

```
E
                                27
                                             36
                     18
VATG ACT GCC GAC AAC AAT AGT ATG CCC CAT GGT GCA GTA TCT AGT TAC GCC AAA
Met Thr Ala Asp Asn Asn Ser Het Pro His Gly Ala Val Ser Ser Tyr Ala Lys
                                                                     18
                                             90
                                                         99
                                 81
                      72
TTA GTG CAA AAC CAA ACA CCT GAA GAC ATT TTG GAA GAG TTT CCT GAA ATT ATT
Leu Val Gln Asn Gln Thr Pro Glu Asp Ile Leu Glu Glu Phe Pro Glu Ile Ile
                                                                     36
                                            144
                     126
                                 135
 CCA TTA CAA CAA AGA CCT AAT ACC CGA TCT AGT GAG ACG TCA AAT GAC GAA AGC
 Pro Leu Gln Gln Arg Pro Asn Thr Arg Ser Ser Glu Thr Ser Asn Asp Glu Ser
                                                                     54
                                                        207
                                                                    216
                                            198
                                189
 GGA GAA ACA TGT TTT TCT GGT CAT GAT GAG GAG CAA ATT AAG TTA ATG AAT GAA
 Gly Glu Thr Cys Phe Ser Gly His Asp Glu Glu Gln Ile Lys Leu Met Asn Glu
                                                                    72
                                             252
                                 243
 AAT TGT ATT GTT TTG GAT TGG GAC GAT AAT GCT ATT GGT GCC GGT ACC AAG AAA
 Asn Cys Ile Val Leu Asp Trp Asp Asn Ala Ile Gly Ala Gly Thr Lys Lys
                                                                    324
                                             306
                                                        315
                                 297
 GTT TGT CAT TTA ATG GAA AAT ATT GAA AAG GGT TTA CTA CAT CGT GCA TTC TCC
 Val Cys His Leu Met Glu Asn Ile Glu Lys Gly Leu Leu His Arg Ala Phe Ser
                                                                    108
                                 351
                                             360
                     342
         333
 GTC TIT ATT TTC AAT GAA CAA GGT GAA TTA CTT TTA CAA CAA AGA GCC ACT GAA
 Val Phe Ile Phe Asn Glu Gln Gly Glu Leu Leu Gln Gln Arg Ala Thr Glu
                                 405
                                             414
                                                         423
                     396
 ARA ATA ACT TTC CCT GAT CTT TGG ACT ARC ACA TGC TGC TCT CAT CCA CTA TGT
 Lys Ile Thr Phe Pro Asp Leu Trp Thr Asn Thr Cys Cys Ser His Pro Leu Cys
                                459
                                                         477
                     450
 ATT GAT GAC GAA TTA GGT TTG AAG GGT AAG CTA GAC GAT AAG ATT AAG GGC GCT
 Ile Asp Asp Glu Leu Gly Leu Lys Gly Lys Leu Asp Asp Lys Ile Lys Gly Ala
                                             522
                                 513
 ATT ACT GCG GCG GTG AGA AAA CTA GAT CAT GAA TTA GGT ATT CCA GAA GAT GAA
 Ile Thr Ala Ala Val Arg Lys Leu Asp His Glu Leu Gly Ile Pro Glu Asp Glu
```

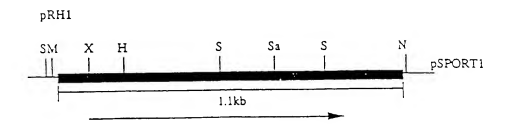
```
558 567 576
                                                   585
ACT AAG ACA AGG GGT AAG TTT CAC TTT TTA AAC AGA ATC CAT TAC ATG GCA CCA
Thr Lys Thr Arg Gly Lys Phe His Phe Leu Asn Arg Ile His Tyr Met Ala Pro
                                         630
                                                    639
                   612
                              621
AGC AAT GAA CCA TGG GGT GAA CAT GAA ATT GAT TAC ATC CTA TTT TAT AAG ATC
Ser Asn Glu Pro Trp Gly Glu His Glu Ile Asp Tyr Ile Leu Phe Tyr Lys Ile
                                         684
                                                    693
                             675
                  666
AAC GCT AAA GAA AAC TTG ACT GTC AAC CCA AAC GTC AAT GAA GTT AGA GAC TTC
Asn Ala Lys Glu Asn Leu Thr Val Asn Pro Asn Val Asn Glu Val Arg Asp Phe
                             729
                                        738
                  720
AAA TGG GTT TCA CCA AAT GAT TTG AAA ACT ATG TTT GCT GAC CCA AGT TAC AAG
Lys Trp Val Ser Pro Asn Asp Leu Lys Thr Met Phe Ala Asp Pro Ser Tyr Lys
                                        792
                  774
                             783
       765
TIT ACG CCT TGG TTT AAG ATT ATT TGC GAG AAT TAC TTA TTC AAC TGG TGG GAG
Phe Thr Pro Trp Phe Lys Ile Ile Cys Glu Asn Tyr Leu Phe Asn Trp Trp Glu
                              837
                                                    855
                   828
CAA TTA GAT GAC CTT TCT GAA GTG GAA AAT GAC AGG CAA ATT CAT AGA ATG CTA
Gln Leu Asp Asp Leu Ser Glu Val Glu Asn Asp Arg Gln Ile His Arg Met Leu
                                                                   F
TAA
```



pHP11

Sp

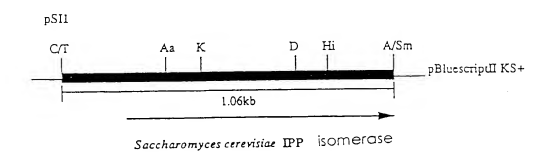
SM



Phaffia rhodozyma IPP isomerase

PPBluescriptII KS+

Haematococcus pluvialis IPP isomerase



Aa: AatII, A: AccII, B:BssHII, D:DraJ, Hi:HincII, H:HpaI, K:KpnI, M:MluI, N:NotI, P:PstI, Sa:SacI, S:SalI, Sp:SphI, X:XbaI

FIG.12

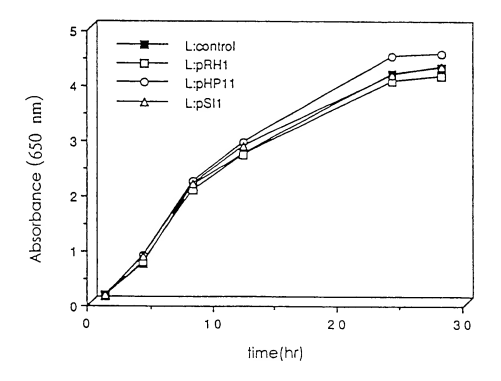


FIG.13

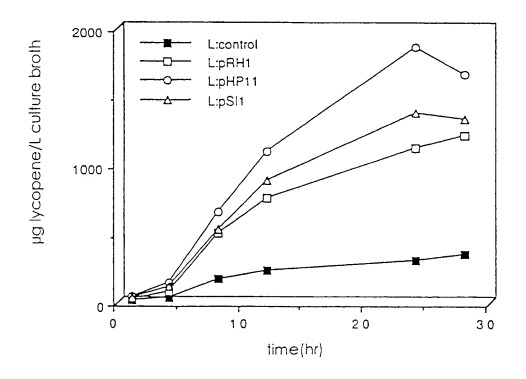


FIG.14

E.coli	µg carotene/g dry weight	production
L: control	228	1
L: pRH1	825	3.6
L: pHP11	1029	4.5
L: pSI1	859	3.8
β: control	488	1
β: pRH1	709	1.5
P: control	246	1
P: pRH1	413	1.7
P: pHP11	504	2.1

INTERNATIONAL SEARCH REPORT

International application No.

PCT/JP96/00574

A. CLASSIFICATION OF SUBJECT MATTER
Int. Cl ⁶ Cl2N15/00, Cl2N9/90
According to International Patent Classification (IPC) or to both national classification and IPC
B. FIELDS SEARCHED
Minimum documentation searched (classification system followed by classification symbols)
Int. C1 ⁶ C12N15/00, C12N9/90
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
BIOSIS PREVIEWS, CAS
C. DOCUMENTS CONSIDERED TO BE RELEVANT
Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No.
A Chu-BIao Xue "A Covalently Constrained Congener of the Saccharomyces cerevisiae Tridecapeptide Mating Pheromone Is an Aronist" J. Biol. Chem., Vol. 264, No. 32, p. 19161-19168
A Ian P. street "Isopentenyldiphosphate: 1 - 4 Dimethylallyldiphosphate Isomerase:Construction of a High-Level Heterologous Expression System for the Gene from Saccharomyces serevisiae and Indentification of an Active-site Nucleophile" Biochemistry, Vol. 29, p. 7351-7538
Further documents are listed in the continuation of Box C. See patent family annex.
* Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance: "E" earlier document but published on or after the international filing date or cannot be considered to earlier the international filing date. "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed. "A" Date of which a count conflict with the application but cited to understand the priority and eritation but cited to understand the priority and eritation but cited to understand the priority date in conflict with the application but cited to understand the priority date in conflict with the application but cited to understand the priority date in conflict with the application but cited to understand the priority and active the international filing date or priority date and not in conflict with the application but cited to understand the priority date and not in conflict with the application but cited to understand the priority claim(s) or which is and not in conflict with the application but cited to understand the priority claim(s) or which is and not in conflict with the application but cited to understand the priority claim(s) or which is and not in conflict with the application but cited to understand the priority date and not in conflict with the application but cited to understand the priority date and not in conflict with the application but cited to understand the priority date and not in conflict with the application but cited to understand the priority date and not in conflict with the application of the inventional filing date or remained. "T" document published after the international filing date or remained and not in conflict with the application but cited t
June 3, 1996 (03. 06. 96) Date of the actual completion of the international search June 11, 1996 (11. 06. 96)
Name and mailing address of the ISA/ Authorized officer
Japanese Patent Office
Facsimile No. Telephone No.

Form PCT/ISA/210 (second sheet) (July 1992)

